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MicroPatent Report

PRODUCTION OF L-PHENYLALANINE BY FERMENTATION

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[57] Abstract:

PURPOSE: To produce L-phenylalanine by a fermentation process in high efficiency by culturing a microorganism transformed with a specific recombinant DNA and collecting the L-phenylalanine produced in the medium. CONSTITUTION: A microorganism is transformed with a recombinant DNA containing a gene coding an enzyme free from feed-back inhibition activity and obtained by substituting an amino acid residue in prephenate dehydratase with other amino acid residue or depleting the amino acid residue. The transformed microorganism is cultured in a medium and the produced L- phenylalanine is separated from the medium. COPYRIGHT: (C)1993,JPO&Japio

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- Recombinant DNA sequences encoding feedback inhibition released enzymes, plasmids comprising the recombinant DNA sequences, transformed microorganisms useful in the production of aromatic amino acids, and a process for preparing aromatic amino acids by fermentation.
- ② A method for releasing feedback inhibition of the key enzymes in the production of aromatic amino acids by termentation is disclosed. Aromatic amino acids are prepared by a process which comprises transforming a microorganism with a recombinant DNA sequence bearing a gene or gene group encoding a feedback inhibition-released enzyme in the phenylalanine and/or tryptophan biosynthetic pathway, obtained by substituting one or two amino acid residue(s) or deleting one or more amino acid residue(s) of 3-deoxy-D-arabinoheptulonic acid 7-phosphate synthase (DS) or prephenate dehydratase, culturing the microorganism and isolating the aromatic amino acid produced in the medium. Higher efficiency and improved yields in the production of L-phenylalanine and L-tryptophan by fermentation are realized.

BACKGROUND OF THE INVENTION

Field of the Invention:

The present invention relates to recombinant DNA sequences encoding feedback inhibition released enzymes, plasmids containing these recombinant DNA sequences, microorganisms transformed with these plasmids, and a process for preparing L-tryptophan, L-phenylalanine and L-tyrosine by fermentation.

Discussion of the Background:

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Demand for aromatic amino acids is rapidly increasing. For example, L-phenylalanine is used as a raw material for the sweetener aspartame, L-tryptophan is an important feed additive, and all three (L-phenylalanine, L-tryptophan, and L-tyrosine) are useful as transfusion drugs.

Many methods for preparing aromatic amino acids using microorganisms are known. For example, methods for preparing L-phenylalanine using recombinant Escherichia coli are described in Japanese Published Unexamined Patent Application Nos. 56-1890, 58-103398, 61-92565 and 1-104160, and World Patent Publication WO 87/00202. A method for preparing L-phenylalanine or L-tyrosine using a mutant belonging to Coryneform bacteria is described in Japanese Patent Published Unexamined Application No. 61-128897, and methods using recombinant Coryneform bacteria are described in Japanese Unexamined Published Patent Application Nos. 60-34197, 60-24192, 61-260892 and 61-124375. A method for preparing L-tryptophan using recombinant E. coli is described in Japanese Published Unexamined Patent Application No. 57-71397 and U.S. Patent No. 4.371,614; methods using mutants of Bacillus subtilis are described in Japanese Published Unexamined Patent Application Nos. 61-104790 and 62-34399; methods using a mutant of Coryneform bacteria are described in Japanese Published Unexamined Patent Application No. 57-174096; and a method using recombinant Coryneform bacteria is described in Japanese Published Unexamined Patent Application No. 57-174096; and a method using recombinant Coryneform bacteria is described in Japanese Published Unexamined Patent Application No. 62-51980.

Generally, in the biosynthetic route of aromatic amino acids, a key enzyme which plays a central role in the biosynthesis is subject to feedback inhibition by the final product. In the methods described above, the desired amino acids are principally produced using microorganisms wherein the key enzyme is released from feedback inhibition by the final product. The key enzymes released from feedback inhibition in the above methods include 3-deoxy-D-arabinoheptulonic acid-7-phosphate synthase (hereafter abbreviated as "DS") and prephenate dehydratase (hereafter abbreviated as "PD").

Turning first to DS, among the microorganisms used in the methods described above. Escherichia coli has three types of naturally-occurring (wild-type) DS isozymes. These isozymes are encoded by genes called aroF, aroG and aroH, which are subject to feedback inhibition by L-tyrosine, L-phenylalanine and L-tryptophan, respectively.

The nucleotide sequences and amino acid sequences relevant to these genes and enzymes are already reported [aroF: Hudson, G.S. and Davidson, B.E., J. Mol. Biol., 180, 1023 (1984); aroG: Davies, W.D. and Davidson, B.E., Nucleic Acids Res., 13, 4045 (1982); aroH: Ray, J.M. et al., J. Bacteriol., 170, 5500 (1988)].

In order to efficiently produce the desired aromatic amino acids, expression of these DS genes must be improved. With respect to aroH-encoded DS, feedback inhibition by L-tryptophan has been released using mutant aroH [Ray, J.M. et al., J. Bacteriol., 170, 5500 (1988)]. However, the DS activity derived from aroH is very poor, and the aroH-derived DS is unsuitable for improvement by recombinant DNA techniques. It is more efficient to utilize aroF- or aroG-encoded DS in which feedback inhibition is released ("feedback inhibition-released" DS).

An example of a mutation which releases feedback inhibition of aroF-encoded DS by L-tyrosine is the substitution of the 148 proline residue from the N-terminus (148 Pro) with a leucine residue [Weaver, L.M. and Herrmann, K.M., J. Bacteriol., 172, 6581 (1980)].

Only a few examples as shown below for the production of aromatic amino acids by fermentation employ feedback inhibition-released DS with a clearly shown mutation site. Edwards et al. teach that feedback inhibition by L-tyrosine in the production of L-phenylalanine by fermentation is released by substituting the 152 glutamine residue (152Gln) of DS encoded by aroF with isoleucine [WO 87/00202]. Furthermore, Sinenki et al. teach that feedback inhibition by L-phenylalanine in the production of L-phenylalanine by fermentation is suppressed by substituting the 76 leucine residue (152Leu) of DS encoded by aroG with valine [Japanese Published Unexamined Patent Application No. 58-103398]. However, the enzyme activity of the feedback inhibition-released DS and the amount of L-phenylalanine produced are unknown. No reports of the production of L-tryptophan by feedback inhibition-released DS mutants are

known.

Turning next to PD, a wild-type bifunctional enzyme (CM-PD) present in Esherichia coli having both chorismate mutase (hereafter abbreviated as "CM") activity and PD activity is subject to feedback inhibition by L-phenylalanine. The enzyme is encoded by a gene called pheA. The nucleotide sequence of pheA and the amino acid sequence of wild-type CM-PD are known [Hudson, G.S. and Davidson, B.E., J. Mol. Biol., 180, 1023 (1984)]. In order to efficiently produce L-phenylalanine, it is important to release the feedback inhibition of CM-PD by L-phenylalanine.

Some examples of modification and mutation on an amino acid level are known to release feedback inhibition for the fermentative production of L-phenylalanine. By modifying two tryptophane residues (226 and 338 amino acids from the N-terminus) of CM-PD with dimethyl(2-hydroxy-5-nitrobenzylsulfonium bromide), an enzyme having resistance to feedback inhibition can be obtained [Gething, M.J.H. and Davidson, B.E., Eur. J. Biochem., 78, 111 (1977)]. Feedback inhibition-released enzyme can be obtained by deleting the 338 tryptophan residue (338 Trp) or substituting 338 Trp and the subsequent residues with arginine-glycine (Japanese Published Unexamined Patent Application No. 1-235597). Inserting the amino acid sequence tryptophan-arginine-serine-proline into the site of the same 338 tryptophan residue also releases feedback inhibition (WO 87/00202). These techniques focus on the 338 tryptophan residue. However, no definitive study on the effects of modifying or mutating the 226 Trp residue has been performed.

On the other hand, in Coryneform bacteria, PD is subject to feedback inhibition by L-phenylalanine. A gene in which the feedback inhibition by L-phenylalanine has been released is known. [Ozaki, A. et al., Agric. Biol. Chem., 49, 2925 (1986); Ito, H. et al., Appl. Microbiol. Biotechnot., 33, 190 (1989)]. The nucleotide sequence of the wild type Coryneform PD gene shows homology to the pheA gene of Escherichia coli K-12 [Follettie, M.T. and Sinsky, A.J., J. Bacteriol., 167, 695 (1986)]. However, the nucleotide sequence of the feedback inhibition-released PD gene in Coryneform bacteria is unknown, as is mutation of the nucleotide sequence and release of feedback inhibition by substitution of the corresponding amino acid sequence.

SUMMARY OF THE INVENTION

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Accordingly, one object of the present invention is to provide a process for efficiently preparing an aromatic amino acids by fermentation.

A further object is to provide transformed microorganisms useful in the production of aromatic amino acids by fermentation.

A further object is to provide recombinant plasmids which express genes encoding key enzymes in the biosynthesis of aromatic amino acids in which feedback inhibition is released.

A further object is to provide recombinant DNA sequences which encode key enzymes in the biosynthesis of aromatic amino acids in which feedback inhibition is released.

A further object is to provide novel recombinant enzymes which are important in the biosynthesis of aromatic amino acids in which feedback inhibition is released.

These and other objects which will become apparent during the following detailed description of the preferred embodiments have been accomplished by a recombinant DNA sequence encoding an enzyme of the aromatic amino acid biosynthetic pathway, wherein feedback inhibition is released, a plasmid comprising the recombinant DNA sequence, a microorganism useful in the production of aromatic amino acids transformed with one or more of the plasmids, and a process for preparing an aromatic amino acid which comprises culturing the transformed microorganism and isolating the aromatic amino acid produced thereby.

BRIEF DESCRIPTION OF THE DRAWINGS

A more complete appreciation of the invention and many of the attendant advantages thereof will be readily obtained as the same becomes better understood by reference to the following detailed description when considered in connection with the accompanying drawings, wherein:

Fig. 1 shows the construction of the plasmids pTS-aroF and pTS-aroG;

Fig. 2 shows the extent of inhibition by L-tyrosine of activity in DS encoded by both wild-type and mutant aroF;

Fig. 3 shows the extent of inhibition by L-phenylalanine of activity in DS encoded by both wild type and mutant aroG.

Fig. 4 shows the extent of inhibition by L-phenylalanine in the prephenate dehydratase activity of both wild-type and mutant chorismate mutase-prephenate dehydratase;

Fig. 5 shows the construction of the plasmid pACKG4.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The present invention relates to a mutant feedback inhibition-released enzyme in the biosynthetic pathway of aromatic amino acids; a recombinant DNA sequence encoding an enzyme of the aromatic amino acid biosynthetic pathway, wherein feedback inhibition is released; a plasmid comprising a recombinant DNA sequence encoding an enzyme of the aromatic amino acid biosynthetic pathway, wherein feedback inhibition is released; a microorganism useful in the production of aromatic amino acids transformed with one or more plasmids comprising a recombinant DNA sequence encoding an enzyme of the aromatic amino acid biosynthetic pathway, wherein feedback inhibition is released; and a process for preparing an aromatic amino acid which comprises culturing a microorganism transformed with one or more plasmids comprising a recombinant DNA sequence encoding an enzyme of the aromatic amino acid biosynthetic pathway, wherein feedback inhibition is released, and isolating the aromatic amino acid produced thereby.

In the present application, the phrase "aromatic amino acid" refers to L-phenylalanine, L-tryptophan and L-tyrosine. Also, an enzyme is "released" from feedback inhibition by a final product if the activity of the enzyme doesn't decrease in the presence of the final product.

Preferably, the enzymes of the present invention in the biosynthetic pathway of aromatic amino acids which are to be released from feedback inhibition are 3-deoxy-D-arabinoheptulonic acid-7-phosphate synthase (DS), prephenate dehydratase (PD) and chorismate mutase-prephenate dehydratase (CM-PD). The means by which each of these enzymes is released from feedback inhibition in the present invention is preferably by mutation, wherein one or two amino acid residue(s) are substituted with other amino acid residue(s) or one or more amino acid residue(s) are deleted. Furthermore, the transformed microorganism preferably belongs to the genus Escherichia, and is preferably transformed with one or more plasmids bearing a recombinant DNA sequence corresponding to one of the above mutant enzymes.

Firstly, the present Inventors have acquired a novel gene encoding DS, wherein the feedback inhibition is released by cloning the natural DS gene of Escherichia coli and subjecting the cloned gene to mutation. Furthermore, the natural PD gene of Brevibacterium lactofermentum is cloned, and a gene encoding PD in which feedback inhibition is released, is cloned from L-phenylalanine-producing Coryneform bacteria. Even further, the natural CM-PD gene of Escherichia coli is cloned, and the cloned gene is then subjected to mutation to produce a novel gene encoding CM-PD in which feedback inhibition is released. By transfecting or transforming phenylalanine-producing bacteria with one or two of the genes of the present invention, production of L-phenylalanine by fermentation can be improved.

The present Inventors have also improved the fermentative production of L-tryptophan by transforming a microorganism with the novel DS gene of the present invention in combination with a tryptophan operon, in which the feedback inhibition Of anthranilate synthase (hereafter abbreviated as AS), an enzyme for the L-tryptophan biosynthesis system, is also released.

The novel gene of the present invention encoding DS having released feedback inhibition is prepared by the following procedure.

Firstly, aroF and aroG genes are cloned from chromosomal DNA of Escherichia coli using the PCR method as described in U.S. Patent Nos. 4,800,159, 4,683,202 and 4,683,195, all incorporated herein by reference. The chromosomal DNA appropriate as a source of the aroF and aroG genes for use in the present invention may be cloned from any strain of Escherichia coli, but the preferred strain is K-12 MC1061 (ATCC 53338). The desired genes are then mutated with hydroxylamine by a known method; for example, that described in J. Mol. Biol., 175, 331 (1984).

The genes aroF and aroG encode DS which is subject to feedback inhibition by L-tyrosine and L-phenylalanine, respectively, and also include mutants caused by genetic polymorphism, etc. Genetic polymorphism refers to a modification of an amino acid sequence of a protein due to natural mutation of a gene.

In order to cause mutation of the gene, a number of effective methods are known. Examples include recombinant PCR methods [PCR Technology, Stockton Press (1989)], site specific mutation [Kramer, W. and Frits, H.J., Methods in Enzymology, 154, 350 (1987)], conventional methods exposing a strain bearing the gene to UV (ultraviolet light) rays, conventional methods treating the DNA or DNA-bearing microorganism with a chemical (N-methyl-N'-nitrosoguanidine, nitric acid, etc.), and conventional methods for chemical synthesis of the desired gene, such as those employing a known automated synthesizer.

In the present invention, the mutated amino acid residue of DS is in the region of the amino acid sequence which participates in the mechanism of feedback inhibition by L-tyrosine, L-phenylalanine or L-tryptophan. For example, in DS encoded by arof, the 147 aspartic acid residue (147 Asp) and the 181 serine

residue (¹⁸ Ser) from the N-terminus are the mutated amino acid residues. Any mutation of the amino acid residue which results in release from feedback inhibition is suitable. For example, substitution, deletion, or addition is suitable. The DS mutations and the corresponding nucleotide sequence mutations exemplified in the present invention are summarized in Table 1.

By transfecting a suitable microorganism with the mutant aroF or aroG gene above as a recombinant DNA sequence, the microorganism can express the recombinant mutant gene in which feedback inhibition is released.

In the present invention, the novel gene encoding PD in Brevibacterium lactofermentum and the novel gene encoding CM-PD in Escherichia coli were prepared as follows.

Firstly, the nucleotide sequence of the Brevibacterium lactofermentum PD gene encoding PD in which feedback inhibition by L-phenylalanine is released was determined and analyzed. It has thus been found that the L-phenylalanine-producing strain expressed PD in which one amino acid is substituted, as compared to the wild strain. Next, based on this finding, a substitution or a deletion of amino acid residue(s) was carried out at the corresponding position of CM-PD in Escherichia coli K-12, resulting in CM-PD in which the feedback inhibition is released.

Enzymes having PD activity referred to in the present invention refer to enzymes derived from microorganisms such as Coryneform bacteria having PD activity, and further refer to enzymes derived from microorganisms such as Escherichia coli, etc., having the bifunctional activity of CM-PD.

In the present invention, the mutated amino acid residue of PD refers to a substitution of an amino acid residue or a deletion of amino acid residue(s) present in the region of the amino acid sequence which participates in the mechanism of feedback inhibition by L-phenylalanine. For example, in PD derived from Brevibacterium lactofermentum, the 235 serine residue (235 Ser) is suitable for mutation, and in CM-PD derived from Escherichia coli, the 330 serine residue (330 Ser) from the N-terminus is an amino acid residue suitable for mutation. Suitable mutations include any which result in the release of feedback inhibition, but particularly suitable mutations include substitutions of 235 Ser or 330 Ser with proline or aspartic acid residue, or deletion of amino acid residues down stream from 330 Ser.

By transfecting a suitable microorganism with the mutant PD or CM-PD gene described above as a recombinant DNA sequence, expression of PD in which feedback inhibition is released is achieved in the transfected microorganism.

The recombinant DNA sequences obtained by the foregoing procedures refer to those obtained by incorporating a gene encoding feedback inhibition-released DS or PD into a vector of plasmid or phage DNA. In the present invention, promoters such as lac, trp, PL, etc. which act in the microorganism may also be used to efficiently perform the expression of the gene. The recombinant DNA sequences referred to herein include those obtained by incorporating one or more of the above-described genes into a chromosome according to known methods. Examples include methods using a transposon (Berg, D.E. and Berg, C.M., Bio/Technol., 1, 417 (1983)), Mu phage (Japanese Published Unexamined Patent Application No. 2-109985) or homologous recombination [Experiments in Molecular Genetics, Cold Spring Harbor Laboratory (1972)].

As the microorganism containing the recombinant DNA, any microorganism may be used, irrespective of species and strain of the microorganism, so long as it expresses the gene encoding the desired enzyme (such as DS or PD) and is capable of producing the aromatic amino acid (for example, in the case of L-phenylalanine, the microorganism which has acquired L-phenylalanine productivity by imparting L-phenylalanine analog resistance thereto). Particularly suitable microorganisms are selected from the genus Escherichia, the genus Brevibacterium, the genus Corynebacterium, the genus Bacillus, the genus Serratia, the genus Pseudomonas, etc.

The thus obtained microorganism transformed by the recombinant DNA bearing the feedback inhibitionreleased DS or PD gene is cultured, the desired aromatic amino acid is produced by the transformed microorganism in a suitable medium, and the accumulated aromatic amino acid is collected and isolated.

The medium used for producing the aromatic amino acid is a conventional medium containing appropriate carbon sources, nitrogen sources, inorganic ions and, if necessary, other organic components.

Suitable carbon sources include sugars such as glucose, lactose, galactose, fructose, starch hydrolysate, etc.; alcohols such as glycerol, sorbitol, etc.; organic acids such as fumaric acid, citric acid, succinic acid, etc.

Suitable nitrogen sources include inorganic ammonium salts such as ammonium sulfate, ammonium chloride, ammonium phosphate, etc.; organic nitrogen such as soybean hydrolysate, etc.; ammonia gas, ammonia water, etc.

Suitable organic trace nutrient sources preferably are present, and include required substances such as vitamin B₁, L-tyrosine, or yeast extract, etc., in an appropriate amount.

In addition thereto, small amounts of potassium phosphate, magnesium sulfate, iron ions, manganese ions, etc. may be present.

Incubation is carried out for 16 to 72 hours under aerobic conditions. The temperature for incubation is maintained between 30 and 45 °C and the pH is maintained in the range of 5 to 7 during the incubation. The pH may be adjusted with either acids or alkaline substances, which may be inorganic or organic, or may be adjusted with ammonia gas, etc., as is appropriate to maintain the desired pH and concentrations of components in the medium.

The desired aromatic amino acid is isolated from the fermentation medium generally by conventional methods, such as use of an appropriate ion exchange resin, precipitation, and/or other known techniques, either alone or in combination.

By the general process described above, the transformant expressing feedback inhibition-released DS, PD and/or CM-PD is obtained, and by culturing the transformant, the productivity of aromatic amino acids can be greatly improved.

Other features of the invention will become apparent in the course of the following descriptions of exemplary embodiments which are given for illustration of the invention, and are not intended to be limiting thereof.

Example 1: Preparation of a novel gene encoding DS in which the feedback inhibition is released

20 (1) Collection of an aroF-derived mutant DS gene of Escherichia coli

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Chromosomal DNA was extracted from Escherichia coli K-12 MC1061 strain in a conventional manner. In a separate procedure, two synthetic DNA primers shown by Sequence Nos. 1 (SEQ ID NO:1) and 2 (SEQ ID NO:2) were synthesized in a conventional manner, based on the known nucleotide sequence of the target aroF gene [J. Mol. Biol., 180, 1023 (1984)].

Sequence No. 1 GCTAACCAGT AAAGCCAACA (SEQ ID NO:1)
Sequence No. 2 CCCACTTCAG CAACCAGTTC (SEQ ID NO:2)

These primers have homologous sequences upstream and downstream from the aroF gene. Using the chromosomal DNA and the DNA primers, PCR (polymerase chain reaction) is conducted according to the method of Erlich et al. [PCR Technology, Stockton Press (1989)], yielding a DNA tragment of 1.5 Kbp. Thereafter, as shown in Fig. 1, left side, the fragment is cleaved with restriction enzymes EcoRV and Eco47III, and the product is then ligated with the Smal digestion product of pHSG398 (manufactured by Takara Shuzo) using T4 DNA ligase. Competent cells of Escherichia coli JM109 strain (manufactured by Takara Shuzo) were transformed with the reaction mixture. A plasmid having the aroF gene was extracted from the strains resistant to chloramphenicol to yield the plasmid pHSG-aroF.

Subsequently, pHSG-aroF was digested with restriction enzymes EcoRI and HindIII, and the resulting DNA fragment bearing the aroF gene was ligated with the EcoRI and HindIII digestion fragment of plasmid pTS1 (Japanese Patent Application No. 2-192162) using T4 DNA ligase. Competent cells of DS-deleted (aroF, aroG, aroH) strain AB3257 of Escherichia coli K-12 were transformed with the reaction mixture (AB3257 strain was acquired from the Escherichia coli Genetic Stock Center). From among the strains resistant to ampicillin, the strain in which auxotrophy of L-tyrosine, L-phenylalanine and L-tryptophan disappeared was selected, and a plasmid was extracted therefrom, yielding plasmid pTS-aroF.

Next, after mutation of plasmid pTS-aroF using hydroxylamine according to the method of J. Mol. Biol., 175, 331 (1984), the mutant was used to transform the E. coli AB3257 strain. After ampicillin-resistant strains were collected, two strains which grew in minimum medium supplemented with 1 mM L-tyrosine were selected. From these strains, plasmids pTS-aroF15 and pTS-aroF33 bearing the genes encoding teedback inhibition-released DS were obtained.

Cells of AB3257 strain transformed with plasmids containing the gene encoding non-feedback inhibition-released DS are subject to feedback inhibition at 1 mM concentration of L-tyrosine in the minimum medium. Accordingly, the strain subject to feedback inhibition failed to synthesize aromatic amino acids such as L-phenylalanine or L-tryptophan, and failed to grow.

(2) Preparation of aroG-derived mutant DS gene of Escherichia coli

A mutant aroG gene was collected in a manner similar to the case of the aroF gene. Two synthetic DNA primers shown by Sequence Nos. 3 (SEQ ID NO:3) and 4 (SEQ ID NO:4) were synthesized in a conventional manner, based on the known nucleotide sequence of the aroG gene (Nucleic Acids Res., 10, 4045 (1982)).

Sequence No. 3 GTATTTACCC CGTTATTGTC (SEQ ID NO:3)
Sequence No. 4 ACTCCGCCGG AAGTGACTAA (SEQ ID NO:4)

Using the primers and the chromosomal DNA of the E. coli MC1061 strain, PCR was carried out to obtain a DNA fragment of 2.1 Kbp. As shown in Fig. 1, right side, the fragment was cleaved with restriction enzymes Sall and Eco47III, and the product was then ligated with the Sall and Smal digestion product of pHSG398 (manufactured by Takara Shuzo) using T4 DNA ligase. Competent cells of Escherichia coli JM109 strain were transformed with the reaction mixture. From among the strains resistant to chloramphenicol, a plasmid having the aroG gene was extracted to yield the plasmid pHSG-aroG.

Subsequently, pHSG-aroG was digested with restriction enzymes EcoRI and Hindlil, and the resulting DNA fragment bearing the aroG gene was ligated with the EcoRI and Hindlil digestion fragment of plasmid pTS1 using T4 DNA ligase. From among the grown strains resistant to ampicillin, the strain in which auxotrophy of L-tyrosine, L-phenylalanine and L-tryptophan disappeared was selected, and a plasmid was extracted therefrom to yield the plasmid pTS-aroG.

Next, after mutation of the plasmid using the hydroxylamine method of Example 1-(1) above, the mutant plasmid was used to transform competent cells of the E. coli AB3257 strain. After ampicillin-resistant strains were isolated, 6 strains which grew in minimum medium supplemented with 10 mM L-phenylalanine were selected. From these strains, plasmids pTS-aroG4, pTS-aroG8, pTS-aroG15, pTS-aroG17, pTS-aroG29 and pTS-aroG40 bearing the aroG gene encoding feedback inhibition-released DS were obtained.

In cells of the AB3257 strain expressing the non-feedback inhibition-released DS, feedback inhibition occurs at a concentration of 10 mM L-phenylalanine in minimum medium. Accordingly, the non-feedback inhibition-suppressed strain fails to synthesize aromatic amino acids such as L-tryptophan and/or L-tyrosine, and therefore, fails to grow.

(3) Determination of DS enzyme activity

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The above plasmids, bearing either mutant aroF (pTS-aroF15 and pTS-aroF33) or mutant aroG (pTS-aroG4, pTS-aroG8, pTS-aroG15, pTS-aroG17, pTS-aroG29 and pTS-aroG40), were used to transform Escherichia coli AB3257 strain having no DS activity. The respective transformants were named AJ 12598 (AB3257/pTS-aroF15), AJ 12599 (AB3257/pTS-aroF33), AJ 12562 (AB3257/pTS-aroG4), AJ 12600 (AB3257/pTS-aroG8), AJ 12563 (AB3257/pTS-aroG15), AJ 12601 (AB3257/pTS-aroG17), AJ 12602 (AB3257/pTS-aroG29) and AJ 12603 (AB3257/pTS-aroG40), respectively. Among them, AJ 12563 and AJ 12603 were deposited as representative strains in the Fermentation Research Institute of the Agency of Industrial Science & Technology of Japan, under the deposit numbers Escherichia coli FERM BP-3567 and FERM BP-3568, respectively. For the purpose of comparison, plasmids hearing wild type genes were also used to transform the E. coli AB3257 strain.

Each of these strains were cultured for 24 hours in a known L-phenylalanine-producing medium [Sugimoto, S. et al., J. Biotechnol., 5, 237 (1988)]. From the culture cells, the crude enzyme solution was prepared by ultrasonic homogenization. The enzyme activity of DS was determined in a conventional manner [Gollub, E. et al., Methods Enzymol., 17, 349], in the presence of L-tyrosine in the case of aroF, and in the presence of L-phenylalanine in the case of aroG. The results presented in Figs. 2 and 3 show that the DS enzyme activity of the wild type transformants (Escherichia coli AB3257/pTS-aroF) is strongly inhibited in the presence of L-tyrosine, whereas the respective mutant transformants are released from feedback inhibition by L-tyrosine. Likewise, in the wild type transformant Escherichia coli AB3257/pTS-aroG, the enzyme activity is strongly inhibited in the presence of L-phenylalanine, whereas in the respective mutant transformants, feedback inhibition by L-phenylalanine is released. Furthermore, the mutant strain AJ 12562 not only releases feedback inhibition by L-phenylalanine, but surprisingly, the DS enzyme activity increases as the concentration of L-phenylalanine increases.

(4) Determination of the mutation site of DS in which the feedback inhibition is released

The nucleotide sequences of the feedback inhibition-released aroF15, aroF33, aroG4, aroG8, aroG15, aroG17, aroG29 and aroG40 were determined in a conventional manner [Molecular Cloning (Second Edition), Cold spring Harbor Press (1989)]. The specific substitution site on the amino acid sequence and the mutation site on the corresponding nucleotide sequence are shown in Table 1.

These sequences are all novel.

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Table 1

Mutant Gene	Substitution	Site of Amino Acid	Corresponding Nucleotide Sequence Change			
	Position from N-Terminus	Amino Acid Sequence Change				
aroF15	147	Asp → Asn	GAT → AAT			
aroF33	181	Ser → Phe	TCC → TTC			
aroG4	150	Pro → Leu	CCA → CTA			
aroG8	202	Ala → Thr	GCC → ACC			
aroG15	146	Asp → Asn	GAT → AAT			
aroG17	147	Met → Ile	ATG → ATA			
	332	Glu → Lys	GAA → AAA			
aroG29	147	Met → Ile	ATG - ATA			
aroG40	157	Met → Ile	ATG → ATA			
	219	Ala → Thr	GCG → ACG			

Example 2: Preparation of a novel gene encoding PD in which the feedback inhibition is released

(1) Determination of the mutation site of Brevibacterium lactofermentum mutant PD

The nucleotide sequence of the Ncol fragment in plasmid pAJ16 bearing the PD gene of Brevibacterium lactofermentum wild strain was determined by the dideoxy method, using the homology to known Corynebacterium s.p. PD gene [Follettie, M.T. and Sinsky, A.J., J. Bacteriot., 167, 695 (1986)] as an index. The plasmid is harbored on Brevibacterium lactofermentum AJ 12125 (FERM P-7546). The resulting nucleotide sequence (SEQ ID NO:5) and the corresponding amino acid sequence (SEQ ID NO:6) are shown below. The B. lactofermentum PD amino acid sequence is different by only one amino acid residue from that of Corynebacterium s.p.

	Sequence 5:
5	ATGAGCGACGCACCAATTGTTGTGGCCTATTTGGGGCCTGCCGGAACCTTCACCGAAGAA MetSerAspAlaProIleValValAlaTyrLeuGlyProAlaGlyThrPheThrGluGlu 1
10	GCCCTCTACAAATTTGCCGACGCCGGCGTATTCGGCGACGGTGAGATCGAGCAGCTACCA AlaLeuTyrLysPheAlaAspAlaGlyValPheGlyAspGlyGluIleGluGlnLeuPro 21 40
	GCCAAATCGCCACAAGAAGCTGTCGACGCGGTCCGCCACGGCACCGCCCAGTTCGCCGTG AlaLysSerProGlnGluAlaValAspAlaValArgHisGlyThrAlaGlnPheAlaVal 41
15	GTCGCCATCGAAAACTTCGTCGACGCCCCGTCACCCCACCTTCGACGCCCTTGACCAG ValAlaIleGluAsnPheValAspGlyProValThrProThrPheAspAlaLeuAspGln 61
20	GGCTCCAACGTGCAAATCATCGCCGAAGAAGAACTCGATATTGCCTTTTCCATCATGGTC GlySerAsnValGlnIleIleAlaGluGluGluLeuAspIleAlaPheSerIleMetVal 81 100
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	ArgProGlyThrSerLeuAlaAspValLysThrLeuAlaThrHisProValGlyTyrGln 101
5	CAAGTGAAAAACTGGATGGCAACCACCATTCCGGACGCCATGTATCTTTCAGCAAGCTCC GlnValLysAsnTrpMetAlaThrThrIleProAspAlaMetTyrLeuSerAlaSerSer
	121 . 140
10	AACGGCGCCGCACAAATGGTTGCCGAAGGAACCGCCGACGCAGCCCAGCGCCCTCC AsnGlyAlaGlyAlaGlnMetValAlaGluGlyThrAlaAspAlaAlaAlaAlaProSer 141
15	CGCGCAGCCGAACTCTTCGGACTGGAACGCCTTGTTGATGATGTCGCCGACGTCCGCGC ArgAlaAlaGluLeuPheGlyLeuGluArgLeuValAspAspValAlaAspValArgGly 181
20	GCCCGCACCCGCTTCGTTGCAGTCCAAGCCCAAGCAGCCGTTTCCGAACCGACCG
	GACCGCACCTCCGTCATTTTCTCCCTACCGAATGTGCCAGGCAGCCTCGTGCGCGCCCTC AspArgThrSerValllePheSerLeuProAsnValProGlySerLeuValArgAlaLeu 201
25	AACGAATTCGCCATCCGTGGCGTCGACCTCACCCGCATCGAATCCCGCCCCACCCGCAAA AsnGluPheAlaIleArgGlyValAspLeuThrArgIleGluSerArgProThrArgLys 221
30	GTCTTCGGAACCTACCGCTTCCACCTGGACATATCCGGACATATCCGCGACATCCCCGTC ValPheGlyThrTyrArgPheHisLeuAspIleSerGlyHisIleArgAspIleProVal 240
35	GCCGAAGCCCTCCGCGCACTCCACCTCCAAGCCGAAGAACTCGTATTCGTCGGTTCCTGG AlaGluAlaLeuArgAlaLeuHisLeuGlnAlaGluGluLeuValPheValGlySerTrp 280
40	CCCTCCAACCGTGCAGAAGACAGCACGCCCCAAACCGACCAACTAGCTAACGTACACAAG ProSerAsnArgAlaGluAspSerThrProGlnThrAspGlnLeuAlaAsnValHisLys 281
	GCGGACGAATGGGTTCGCGCAGCAAGCGAAGGAAAGTTAACTAG AlaaspGluTrpValargalaalaSerGluGlyArgLysLeuAsn*** 301
4 5	

Next, the nucleotide sequence (SEQ ID NO: 7) of the gene on plasmid pPH14 encoding PD of the phenylalanine-producing strain of Brevibacterium lactofermentum was determined. The sequence shown below was obtained. The plasmid used was the one borne on Brevibacterium lactofermentum AJ 12259 (FERM BP-3565). A comparison of the amino acid sequences was made between the wild-type PD and the feedback inhibition-released PD (SEQ ID NO: 8) and it was found that ²³⁵ Ser residue of the wild strain was mutated to a proline residue in the feedback inhibition-released PD.

	Sequence of <u>Brevibacterium</u> <u>lactofermentum</u> PD (pPH14):
5	ATGAGCGACGCACCAATTGTTGTGGCCTATTTGGGGCCTGCCGGAACCTTCACCGAAGAI MetSerAspAlaProlleValValAlaTyrLeuGlyProAlaGlyThrPheThrGluGluGluGlyThrPheThrGluGluGluGluGluGluGluGluGluGluGluGluGluG
o	GCCCTCTACAAATTTGCCGACGCCGGCGTATTCGGCGACGGTGAGATCGAGCAGCTACCAAAATTTGCCGACGCGCGCG
	GCCAAATCGCCACAAGAAGCTGTCGACGCGGTCCGCCACGGCACCGCCCAGTTCGCGGTCAAAAAAAA
5	GTCGCCATCGAAAACTTCGTCGACGGCCCCGTCACCCCCACCTTCGACGCCCCTTGACCAC ValAlaIleGluAsnPheValAspGlyProValThrProThrPheAspAlaLeuAspGlr 61
o	GGCTCCAACGTGCAAATCATCGCCGAAGAAGAACTCGATATTGCCTTTTCCATCATGGTCGlySerAsnValGlnIleIleAlaGluGluGluLeuAspIleAlaPheSerIleMetVal
5	CGGCCAGGGACTTCGCTTGCCGACGTCAAAACCCTCGCCACCCAC
0	CAAGTGAAAAACTGGATGGCAACCACCATTCCGGACGCCATGTATCTTTCAGCAAGCTCCGInValLysAsnTrpMetAlaThrThrIleProAspAlaMetTyrLeuSerAlaSerSer 121
	AACGGCGCCGCGCACAAATGGTTGCCGAAGGAACCGCCGACGCAGCGCAGCGCCCTCCASnGlyAlaGlyAlaGlnMetValAlaGluGlyThrAlaAspAlaAlaAlaAlaProSer
5	CGCGCAGCCGAACTCTTCGGACTGGAACGCCTTGTTGATGATGTCGCCGACGTCCGCGGCACGTCCGCGGCACGTCCGCGGCGACGTCCGCGGCGACGTCCGCGGCGACGTCCGCGGCGACGTCCGCGGCGACGTCCGCGGCGACGTCCGCGGCGACGTCCGCGGACGTCCGCGGACGTCCGCGGACGTCCGCGGACGTCCGCGGACGTCCGCGGACGTCGACGTCGCGACGTCGCGACGTCGCGACGTCGCGACGTCGCGACGTCGCGACGTCGCGACGTCGAACGTCGAACGTCGAACGTCGAACGTCGAACGTCGAACGTCGAACGTCGAACGTCGAACGTCGAACGTCGCGACGTCGAACGTCGAACGTCGAACGTCGAACGTCAACAACAACAACAACAACAACAACAACAACAACAACAA
0	

AlaArgThrArgPheValAlaValGlnAlaGlnAlaAlaValSerGluProThrGlyHis 5 GACCGCACCTCCGTCATTTTCTCCCTACCGAATGTGCCAGGCAGCCTCGTGCGCGCCCTC AspArgThrSerValllePheSerLeuProAsnValProGlySerLeuValArgAlaLeu 10 AsnGluPheAlaIleArgGlyValAspLeuThrArgIleGluProArgProThrArgLys **GTCTTCGGAACCTACCGCTTCCACCTGGACATATCCGGACATATCCGCGACATCCCCGTC** ValPheGlyThrTyrArgPheHisLeuAspIleSerGlyHisIleArgAspIleProVal 15 GCCGAAGCCCTCCGCGCACTCCAAGCCGAAGAACTCGTATTCGTCGGTTCCTGG AlaGluAlaLeuArgAlaLeuHisLeuGlnAlaGluGluLeuValPheValGlySerTrp CCCTCCAACCGTGCAGAAGACAGCACGCCCCAAACCGACCAACTAGCTAACGTACACAAG ProSerAsnArgAlaGluAspSerThrProGlnThrAspGlnLeuAlaAsnValHisLys 25 GCGGACGAATGGGTTCGCGCAGCAAGCGAAGGAAGGAACTTAACTAG AlaAspGluTrpValArgAlaAlaSerGluGlyArgLysLeuAsn*** 315 (2) Construction of a novel gene encoding a mutant CM-PD of Escherichia coli Chromosomal DNA was extracted from Escherichia coli K-12 RR1 strain in a conventional manner. In a separate procedure, four synthetic DNA primers (Sequence Nos. 7-10) (SEQ ID NOS: 9-12) were 35 chemically synthesized in a conventional manner, based on the known nucleotide sequence of the pheA gene [Hudson, G.S. and Davidson, B.E., J. Mol. Biol., 180, 1023 (1984)]. Sequence No. 7 TCAACAAGCT GGAACGGACG (SEO ID NO: 9) Sequence No. 8 CGCCGATTTA CCGCCTTGAG (SEQ ID NO: 10) 45 Sequence No. 9 CCGTCTGGAA CCACGCCCGA T (SEQ ID NO: 11) Sequence No. 10 ATCGGGCGTG ATTCCAGACG G (SEQ ID NO: 12) Sequence Nos. 7 and 8 have homologous sequences upstream and downstream from the pheA gene, respectively. Sequence Nos. 9 and 10 are complementary to each other and have almost perfect homology to the sequence around the 330 serine residue (330 Ser), except that one base pair is different, i.e., T (thymine base) is substituted with C (cytosine base). CM-PD of Escherichia coli K-12 has high homology to

becomes a proline residue.

PD of Brevibacterium lactofermentum. In particular, the 330 serine residue from the N-terminal (330 Ser) of the CM-PD of Escherichia coli K-12 corresponds to the 235 serine residue (235 Ser) of Brevibacterium lactofermentum PD. Sequence Nos. 9 and 10 are synthesized in such a manner that the 330 serine residue

Next, using 1 µg of the chromosomal DNA and either 300 ng of each of the primers of Sequence Nos. 1 and 10, or 300 ng of each of the primers of Sequence Nos. 8 and 9, PCR was carried out to obtain DNA fragments of 1.3 Kbp and 0.5 Kbp, respectively. The PCR temperature cycle of reaction at 94 °C for one minute, at 50 °C for 2 minutes and at 72 °C for 3 minutes was repeated for 20 cycles using a continuous replication reaction device (Thermal Cycler, manufactured by Perkin Elmer Cetus Co.), according to the method of Erlich et al. [PCR Technology, Stockton Press (1989)]. These DNA fragments were subjected to agarose gel electrophoresis and recovered using a standard DNA recovery kit (Gene Clean, manufactured by Funakoshi Co.).

Separately, using these fragments and the primers of Sequence Nos. 7 and 8, PCR reaction was further carried out to obtain a DNA fragment of 1.8 Kbp. After the 1.8 Kbp fragment was digested with BamHI and PstI, a DNA fragment of 1.7 Kbp was recovered by agarose electrophoresis. Subsequently, the 1.7 Kbp fragment was ligated with the BamHI and PstI digestion product of plasmid pHSG398 (manufactured by Takara Shuzo) using T4 DNA ligase. The ligation product was used to transfect Escherichia coli KA197 strain (pheA). Among the strains resistant to chloramphenicol, the strain in which phenylalanine auxotrophy disappeared was selected, and a plasmid was recovered. The plasmid was named pPHAB. Its nucleotide sequence was determined. This plasmid bears the mutant CM-PD enzyme gene in which the 330 serine residue was substituted with a proline residue.

Also using the same methods mentioned above, 330 serine residue from the N-terminal (³³⁰Ser) of the CM-PD of Escherichia coli K-12 was substituted with an aspartic acid residue. Sequence Nos. 11 and 12 were synthesized in such a manner that the 330 serine residue became an aspartic acid residue.

Nos. 11 COGTCTGGAA GACCGCCCGA T

Nos. 12 ATCGGGCGGT CTTCCAGACG G

In this way, the plasmid pPHAD, which bears the mutant CM-PD enzyme gene in which the 330 serine residue was substituted with an aspartic acid, was obtained.

Also using the same methods mentioned above, amino acid residues downstream from ³³⁰Ser of the CM-PD of Escherichia coli K-12 were deleted. Sequence Nos. 13 and 14 were synthesized in such a manner that the codon of the 330 serine residue became termination codon.

Nos. 13 CCGTCTGGAA TGACGCCCGA T

Nos. 14 ATCGGGCGTC ATTCCAGACG G

In this way, the plasmid pPHATerm, which bears the mutant CM-PD enzyme gene in which the amino acid residues downstream from the 330 serine residue were deleted, was obtained.

(3) Construction of a tyrA gene-defected W3110 strain of Escherichia coli K-12

Escherichia coli W3110 strain (acquired from National Institute of Heredity) was spread on a plate medium containing streptomycin to obtain a streptomycin-resistant strain. Next, this strain was cultured in a culture medium in combination with Escherichia coli K-12 ME8424 strain (HfrPO45, thi, relA1, tyrA: : Tn10, ung-1, nadB) (acquired from National Institute of Heredity), and the medium was allowed to stand at 37 °C for 15 minutes to perform conjugation transfer. Then the medium was applied to a plate medium containing streptomycin, tetracycline and L-tyrosine. The formed colony, i.e., Escherichia coli W3110 (tyrA) strain, was collected.

The plasmid pPHAB, pPHAD, and pPHATerm obtained in Example (2)-2 above were used to transform competent cells of the E. coli W3110 (tyrA) strain. The transformant Escherichia coli K-12 [W3110 (tyrA)/pPHAB] Escherichia coli K-12 [W3110 (tyrA)/pPHAD], and Escherichia coli K-12 [W3110 (tyrA)/pPHATerm] were deposited in the Fermentation Research Institute of the Agency of Industrial Science & Technology of Japan. The deposit numbers are FERM BP-3566, FERM BP-12659, and FERM BP-12662, respectively.

(4) Measurement of PD enzyme activity

Escherichia coli K-12 W3110 (tyrA)/(pPHAB) strain was cultured at 37°C for 15 hours in L medium and

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the cells were collected by centrifugation. Then, the cells were washed twice with physiological saline, and suspended in 250 mM Tris-hydrochloride buffer (pH 7.5) containing 0.5 mM dithiothreitol under ice cooling. By ultrasonication (20 KHz) for 30 seconds four times, the crude enzyme solution was prepared.

The PD enzyme activity was determined in a conventional manner [cotton, R.G.H. and Gibson, F., Meth. in Enzymol., 17, 564 (1970)]. Using the crude enzyme, the enzymatic reaction was carried out at 37°C for 10 minutes in the presence of 50 mM Tris-hydrochloride buffer (pH 8.2) containing 1 mM barium prephenate and 0.5 mM L-tyrosine. Aqueous sodium hydroxide (1 N) was added to terminate the reaction, and the formed phenylpyruvic acid was measured at an extinction wavelength of 320 nm.

Quantitative determination of protein was made using the Protein Assay Kit (manufactured by Bio Rad Co.), according to the protocol of the manufacturer.

The results presented in Fig. 4 show that the enzyme reaction in strains transformed with the wild CM-PD gene was strongly inhibited in the presence of 0.5 mM L-phenylalanine, whereas strains transformed with a mutant CM-PD gene exhibited almost no inhibition, even in the presence of 5 mM L-phenylalanine.

Further, in the case of the plasmid bearing the wild type enzyme gene, in the absence of L-phenylalanine, the enzyme activity was only 3.5 x 10² U/mg protein. In the case of the mutant CM-PD gene, the activity was 1.5 x 10⁴ U/mg protein. The results show that not only is expression of the mutant type CM-PD enzyme gene released from feedback inhibition by L-phenylalanine in transformants containing the mutant gene, but surprisingly, the amount of enzyme and/or enzyme activity can be increased by roughly two orders of magnitude.

By using the same method above, the PD enzyme activities of Escherichia coli W3110 (tyrA)/(pPHAD) and Escherichia coli W3110 (tyrA)/(pPHATerm) were determined. As the result, the PD enzymes of the both strains were found to be released from feedback inhibition by L-phenylalanine.

Example 3: Production of L-phenylalanine by fermentation

(1) Construction of Escherichia coli K-12 bearing a CM-PD gene alone and in combination with a DS gene, in which the feedback inhibition is released

From pTS-aroG4 bearing the feedback inhibition-released DS gene obtained in Example 1, the aroG4 portion was excised with restriction enzymes EcoRI and HindIII. The fragment was inserted into the cleavage site of pBR322 with EcoRI and HindIII to obtain plasmid pBR-aroG4 (having an ampicillin-resistant marker).

In a separate procedure, pPHAB bearing the feedback inhibition-released CM-PD gene obtained in Example 2 was digested with restriction enzymes BamHI and HindIII to excise the fragment containing the CM-PD gene. This fragment was inserted into the cleavage site of pACYC184 with BamHI and HindIII to construct plasmid pACMAB (selection marker was chloramphenicol resistance). The pACAMB plasmid was used to transform competent cells of Escherichia coli K-12 W3110 (tyrA), yielding transformant W3110 (tyrA)/pACMAB.

Furthermore, the two plasmids pACMAB and pBR-aroG4 were used to transform W3110 (tyrA) yielding transformant W3110 (tyrA)/pBR-aroG4,pACMAB. The transformant W3110 (tyrA)/pBR-aroG4,pACMAB was named AJ 12604 strain and deposited in Fermentation Research Institute of the Agency of Industrial Science & Technology of Japan under the deposit number FERM BP-3579.

(3) Production of L-phenylalanine

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The transformant AJ 12604, W3110 (tyrA)/pACMAB, W3110 (tyrA)/pPHAD, W3110 (tyrA)/pPHATerm, and W3110 (tyrA) described above were cultured at 37°C for 24 hours in L-phenylalanine-producing medium (containing 20 g of glucose, 29.4 g of disodium hydrogen phosphate, 6 g of potassium dihydrogen phosphate, 1 g of sodium chloride, 2 g of ammonium chloride, 10 g of sodium citrate, 0.4 g of sodium glutamate, 3 g of magnesium sulfate heptahydrate, 0.23 g of calcium chloride and 2 mg of thiamine hydrochloride in 1 liter of water). The results are shown in Table 2. Quantitative assay was performed by high performance liquid chromatography. An outstanding improvement in the fermentative production of L-phenylalanine using the AJ 12604 strain was obtained.

Table 2

Strain	Amount of L-phenylalanine
W3110(tyrA)	0.1
W3110 (tyrA)/pACMAB	0.5
W3110 (tyrA)/pPHAD	0.5
W3110 (tyrA)/pPHATerm	0.5
AJ 12604	3.8
AU 12004	3.0

Example 4: Production of L-tryptophan by fermentation

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(1) Construction of a plasmid bearing feedback inhibition-released DS

Plasmid pACYC177 (acquired from National Institute of Heredity; ampicillin resistance, 3.6 Kbp) was digested with restriction enzyme Xhol. After the digestion site was made blunt ended by Klenow treatment, an EcoRI linker was ligated therewith using T4 DNA ligase to obtain the plasmid pACYC177E, in which the Xhol site became EcoRI. Next, the plasmid pTS-aroG4 described in Example 1-(2) and 1-(3) above was digested with restriction enzymes EcoRI and HindIII to obtain the fragment containing aroG4. This fragment was ligated with the EcoRI- and HindIII-digested pACYC177E using T4 DNA ligase. Competent cells of the AB3257 strain (described in Example 1) was transformed with the reaction mixture. Among the ampicillin-resistant strains grown, a strain in which auxotrophy for each of L-tyrosine, L-phenylalanine and L-tryptophane disappeared was selected, and a plasmid was extracted. Thus, plasmid pACEG4 (5.1 Kbp) was obtained. An EcoRI-EcoRI fragment containing a gene conferring kanamycin resistance (Kanamycin Gene Block; 1.3 Kbp, manufactured by Pharmacia Fine chemicals) was ligated with the plasmid pACEG4 at the EcoRI site using T4 DNA ligase, thus yielding the plasmid pACKG4 (resistant to ampicillin and kanamycin, 6.4 KbP). The procedure of the construction of pACKG4 is outlined in Fig. 5.

(2) Construction of Escherichia coli K-12 bearing a feedback inhibition-released DS gene and a tryptophan operon

Competent cells of Escherichia coli K-12 AGX6aroP strain (described in U.S. Patent No. 4,371,614, incorporated herein by reference; deposit number: NRRL B-12264) bearing the plasmid pGX50 harboring a tryptophan operon was transformed with the pACKG4 plasmid described above to obtain Escherichia coli AGX6aroP/pGX50,pACKG4. The genotype of Escherichia coli AGX6aroP strain is tna, trpR+, aroP.

(3) Production of L-tryptophan

The transformant Escherichia coli AGX6aroP/pGX50,pACKG4 and AGX6aroP/pGX50 described above was cultured at 30 °C for 72 hours in L-tryptophan-producing medium (containing 40 g of glucose, 15 g of ammonium sulfate, 1 g of potassium monohydrogen phosphate, 1 g of magnesium sulfate heptahydrate, 0.01 g of ferrous sulfate heptahydrate, 0.01 g of manganese chloride tetrahydrate, 2 g of yeast extract and 40 g of calcium carbonate in 1 liter of water, pH 7). The results are shown in Table 3. Quantitative assay of L-tryptophan was performed by high performance liquid chromatography. An outstanding improvement in the fermentative production of L-tryptophan using the AGXaroP/pGX50,pACKG strain was obtained.

Table 3

Strain	Amount of L-tryptophan Produced (g/l)
AGX6aroP/pGX50	0.15
AGX6aroP/pGX50,pACKG4	0.45

Obviously, numerous modifications and variations of the present invention are possible in light of the

above teachings. It is therefore to be understood that within the scope of the appended claims, the invention may be practiced otherwise than as specifically described herein.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

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(i) APPLICANT: Ajinomoto Co., Ltd.

(ii) Title of invention: Genes encoding feedback inhibition-released enzymes, plasmids containing the genes, and microorganisms transformed with the plasmids USEFUL IN PROCESSES FOR PREPARING AROMATIC AMINO ACIDS BY 15

(iii) NUMBER OF SEQUENCES: 12

PERMENTAT

(2) INFORMATION FOR SEQ ID NO:1:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 20 base pairs

(B) TYPE: nucleic acid

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		(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
5	(iii)	HYPOTHETICAL: NO	
10	(vi)	ORIGINAL SOURCE: (A) ORGANISM: Escherichia coli (B) STRAIN: K-12 MC1061	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:1:	
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	(2) INFO	RMATION FOR SEQ ID NO:2:	
20	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
25	(iii)	HYPOTHETICAL: NO	
	(vi)	ORIGINAL SOURCE: (A) ORGANISM: Escherichia coli (B) STRAIN: K-12 HC1061	
30	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:2:	
	•	AG CAACCAGTTC	20
35	(2) INPO	RMATION FOR SEQ ID NO:3:	
40	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDWESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
45	(iii)	HYPOTHETICAL: NO	

	(vi) ORIGINAL SOURCE: (A) ORGANISM: Escherichia coli (B) STRAIN: K-12 MC1061	
5	, ,	
,	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
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	(2) INFORMATION FOR SEQ ID NO:4:	
15	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	.•
	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOTHETICAL: NO	
20		
	(vi) ORIGINAL SOURCE: (A) ORGANISM: Escherichia coli (B) STRAIM: K-12 MC1061	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
	ACTCCGCCGG AAGTGACTAA	20
		••
30	(2) INFORMATION FOR SEQ ID NO:5:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 948 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOTHETICAL: NO	
40	(vi) ORIGINAL SOURCE: (A) ORGANISM: Brevibacterium lactofermentum (B) STRAIN: AJ 12125	
45	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1945	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

		,	,					-									
5	ATG Met 1	AGC Ser) Asp	CCA Ala	CCA Pro 5	ATT	GTT Val	GTG Val	GCC Ala	TAT Tyr 10	Lev	GGG (Gl)	ccr Pro	GCC Ala	GGA Gly 15	Thr	48
		ACC Thr								Ala					. Phe	GGC Gly	96
10		GGT Gly							Ala					Glu		GTC Val	144
15		GCG Ala 50						Ala					. Val			GAA Glu	192
		TTC Phe										Asp				CAG Gln 80	240
20		TCC Ser									Glu					Phe	288
25		ATC Ile													Thr	CTC Leu	336
		ACC												Met		ACC Thr	384
30		ATT Ile 130											λεπ			GGC Gly	432
35		CAA Gln															480
		GCA Ala														GCC Ala	528
40		GTC Val															576
45		GTT Val															624

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	CTA Leu	CCG Pro 210	Asn	GTG Val	CCA Pro	eja Gec	AGC Ser 215	Leu	GTG Val	CGC Arg	GCC	CTC Leu 220	Asn	GAA Glu	TTC Pbe	GCC Ala	672
5							Thr					Arg				AAA Lys 240	720
10	GTC Val	TTC Phe	GCA	ACC Thr	TAC Tyr 245	CGC Arg	TTC Phe	CAC His	CTG Leu	GAC Asp 250	Ile	TCC Ser	GGA Gly	CAT His	ATC 11e 255	Arg	768
	gac Asp	ATC Ile	CCC Pro	GTC Val 260	GCC Ala	GAA Glu	GCC Ala	cTC Leu	CGC Arg 265	λla	CTC Leu	CAC His	CTC Leu	CAA Gln 270	Ala	GAA Glu	816
15	GAA Glu	CTC Leu	GTA Val 275	TTC Phe	GTC Val	GGT Gly	TCC Ser	TGG Trp 280	CCC Pro	TCC Ser	AAC A≤n	CGT Arg	GCA Ala 285	Glu	GAC Asp	AGC Ser	864
20		CCC Pro 290											λla			TGG Trp	912
au	GTT Val 305	CGC Arg	GCA Ala	GCA Ala	AGC Ser	GAA Glu 310	GGA Gly	AGG Arg	AAA Lys	CTT Leu	AAC Asn 315						948
25	(2)	INPO	ORMA?	MOI	FOR	eeq	ID P	10:6	:								
		1	(i) £	(A)	ENCE TYI TOI	GTH:	315	an	ino a id		•						
30			(i))							. Th	WO .	• •					
35	Met 1	Ser											Pro	Ala	Gly 15	Thr	
	Phe	Thr	Glu	Glu 20	Ala	Leu	Tyr	Lys	Phe 25	Ala	qaA	Ala	Gly	Val 30	Phe	Gly	
40	Asp	Gly	G] \u 35	Ile	Glu	Gln	Leu	Pro 40	Ala	Lys	Ser	Pro	Gln 45	Glu	Ala	Val	
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	Asn 65		Val	dsv	Gly	Pro 70	Val	Thr	Pro	Thr	Phe 75	Asp	Ala	Leu	Asp	Glr 80
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	Ser	Ile		Val 100	Arg	Pro	Gly	Thr	Ser 105	Leu	Ala	Asp	Val	Lys 110	Thr	Leu
0	Ala	Thr	His 115	Pro	Val	Gly	Tyr	Gln 120	Gln	Val	Lys	neA	Trp 125	Met	Ala	The
5	Thr	Ile 130	Pro	yeb	Ala	Het	Tyr 135	Lou	Ser	Ala	Ser	Ser 140	Asn	Gly	Ala	Gly
	Ala 145	Gln	Net	Val	Ala	Glu 150		Thr		Asp	Ala 155	Ala	Ala	Ala	Pro	Ser 160
0	Arg	Ala	Ala	Glu	Leu 165	Phe	Gly	Leu	Glu	Arg 170	Leu	Val	Asp	Asp	Val 175	Ala
	Хsр	Val	Arg	Gly 180	Ala	λrg	Thr	Arg	Phe 185		Ala	Val	Gln	Ala 190	Gln	Ala
5	Ala	Val	Ser 195	Glu	Pro	Thr	Gly	His 300	Asp	Arg	Thr	Ser	Val 305	Ile	Phe:	Ser
	Leu	Pro 210	Asn	Val	Pro	Gly	Ser 215	Leu	Val	Arg	Ala	Leu 220	Asn	Glu	Phe	Ala ·
o	11e 225	Arg	Gly	Val	Хsр	Leu 230	Thr	Arg:	Ile	Glu	Ser 235	Arg	Pro 	Thr	Arg	Lys 240
	Val	Phe	Gly	Thr	Tyr 245	Arg	Phe	His	Lou	Asp 250	Ile	Ser	Gly	His	Ile 355.	Arg
5	qaA	Ile	Pro	Val 260	Ala		Ala		Arg 265	Ala	Leu	His.		Gln 270	Ala	Glu
o	Glu	Leu	Val 275	Phe	Val	Gly	Ser	Trp	Pro	Ser	Asn _.		Ala 285	Glu	qsA	Ser
	Thr	Pro 290	Gln	Thr	Asp	Gln	Leu 295	Ala	Asn-	Val	His	Lys 300	Ala	qzƙ	Glu	Trp
5	Val 305	Arg	Ala	Ala	Ser	G1u 310	Gly	Arg	Lys		Asn 315					
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(2) INFORMATION FOR SEQ ID NO:7:

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(1) SEQUENCE CHARACTERISTICS:

	(A) LENGTH: 948 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
5	(ii) MOTECULE TYPE: DNA (genomia)	
	(111) HYPOTHETICAL: NO	
10	(vi) ORIGINAL SOURCE:(A) ORGANISM: Brevibacterium lactofermentum(B) STRAIN: AJ 12259	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
15	ATGAGCGACGCACCAATTGTTGTGGCCTATTTGGGGCCTGCCGGAACCTTCACCGAAGAA	60
	GCCCTCTACAAATTTGCCGACGCCGGCGTATTCGGCGACGGTGAGATCGAGCAGCTACCA	120
	GCCAAATCGCCACAAGAAGCTGTCGACGCGGTCCGCCACGGCACCGCCCAGTTCGCGGTG	180
20	GTCGCCATCGAAAACTTCGTCGACGGCCCCGTCACCCCACCTTCGACGCCCTTGACCAG	240
	GGCTCCAACGTGCAAATCATCGCCGAAGAAGAACTCGATATTGCCTTTTCCATCATGGTC	300
	CGGCCAGGGACTTCGCTTGCCGACGTCAAAACCCTCGCCACCCAC	360
25	CAAGTGAAAACTGGATGGCAACCACCATTCCGGACGCCATGTATCTTTCAGCAAGCTCC	420
	AACGGCGCCGCACAAATGGTTGCCGAAGGAACCGCCGACGCAGCCGCAGCGCCCTCC	480
	CGCGCAGCCGAACTCTTCGGACTGGAACGCCTTGTTGATGATGTCGCCGACGTCCGCGGC	540
30	GCCCGCACCCGCTTCGTTGCAGTCCAAGCCCAAGCAGCCGTTTCCGAACCGACCG	600
	GACCGCACCTCCGTCATTTTCTCCCTACCGAATGTGCCAGGCAGCCTCGTCCGCGCCCTC	660
	AACGAATTCGCCATCCGTGGCGTCGACCTCACCGGCATCGAATCCCGCCCCACCCGCAAA	720
35	GTCTTCGGAACCTACCGCTTCCACCTGGACATATCCGGACATATCCGCGACATCCCCGTC	780
	GCCGAAGCCCTCCGCGCACTCCACCTCCAAGCCGAAGAACTCGTATTCGTCGGTTCCTGG	840
	CCCTCCAACCGTGCAGAAGACAGCCCCCCAAACCGACCAACTAGCTAACGTACACAAG	900
40	GCGGACGAATGGGTTCGCGCAGCAAGCGAAGGAAACTTAACTAG	948
	(2) INFORMATION FOR SEQ ID NO:8:	

(i) SEQUENCE CHARACTERISTICS:

(A)	Length:	315	amino	acids
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- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SPQUENCE DESCRIPTION: SEQ ID NO.81

- Met Ser Asp Ala Pro Ile Val Val Ala Tyr Leu Gly Pro Ala Gly Thr
 1 5 10 15
 - Phe Thr Glu Glu Ala Leu Tyr Lys Phe Ala Asp Ala Gly Val Phe Gly 20 25 30
- Asp Gly Glu Ile Glu Gln Leu Pro Ala Lys Ser Pro Gln Glu Ala Val
 - Asp Ala Val Arg His Gly Thr Ala Gln Phc Ala Val Val Ala Ile Glu
 50 55 60
 - Asn Phe Val Asp Gly Pro Val Thr Pro Thr Phe Asp Ala Leu Asp Gln 65 70 75 80
 - Gly Ser Asn Val Gln Ile Ile Ala Glu Glu Glu Leu Asp Ile Ala Phe 85 90 95
 - Ser Ile Met Val Arg Pro Gly Thr Ser Leu Ala Asp Val Lys Thr Leu
 100 105 110
- Ala Thr His Pro Val Gly Tyr Gln Gln Val Lys Asn Trp Met Ala Thr
 120 125
 - Thr Ile Pro Asp Ala Met Tyr Leu Ser Ala Ser Ser Asn Gly Ala Gly 130 140
- Ala Gln Met Val Ala Glu Gly Thr Ala Asp Ala Ala Ala Ala Pro Ser 145 150 155 160
 - Arg Ala Ala Glu Leu Phe Gly Leu Glu Arg Leu Val Asp Asp Val Ala 165 170 175
- 40 App Val Arg Gly Ala Arg Thr Arg Phe Val Ala Val Gln Ala Gln Ala 180 185 190
 - Ala Val Ser Glu Pro Thr Gly His Asp Arg Thr Ser Val Ile Phe Ser 195 200 205
- Lou Pro Asn Val Pro Gly Ser Leu Val Arg Ala Leu Asn Glu Phe Ala 210 215 220
- Ile Arg Gly Val Asp Leu Thr Arg Ile Glu Pro Arg Pro Thr Arg Lys 225 230 235 240

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	Va)	Phe Gly	Thr Tyr		Phe	His	Leu	հեր 250	Ile	Ser	Gly	His	Ila 255	Arg	
5	Asy	lle Pro	Val Ala 260	Glu	λla	Leu	Arg 265	Ala	Leu	His	Leu	Gln 270	Ala	Glu	
	Gli	Leu Val 275	Phe Val	Gly	Ser	Trp 280	Pro	Ser	Asn	Arg	Ala 285	Glu	Аєр	Ser	
10	Thr	Pro Gln 290	Thr Asp	Gln	Leu 295	λla	Asn	Val	His	Lys 300	λla	Asp	Glu	Trp	
	Va.) 305	Arg Ala	Ala Ser	Glu 310	Gly	Arg	Lys	Leu	As n 315						
15	(2) INFO	RMATION :	FOR SEQ	ID NO):9:										
20	(i)	(B) TYI (C) STI	E CHARAC NGTH: 20 PE: nucl RANDEDHE POLOGY:	base eic a SS: s	pai icid ingl	irs									
	(11)	MOLECULI	E TYPE:	DNA (geno	mic)									
	(iii)	HYPOTHE	FICAL: N	0											
25	1 4.4	OPICINA	COMPON	_											
	(41)	ORIGINAL (A) OR	GANISM:		rich	da c	oli								
•			RAIN: K-												
30	(xi)	SEQUENCI	e descri	PTION	l: SE	Q ID	NO:	9:							
	TCAACAAG	CT GGAACO	GACG												20
35	(2) INFO	RMATION I	POR SEQ	ID NO	:10:										
40	(1)	(B) TYP (C) STR	CHARAC NGTH: 20 PE: nucl RANDEDNE POLOGY:	basa eic a SS: s	pai cid ingl	rs									
	(ii)	MOLECULE				mic)									
	(iii)	нурогнет	'ICAL: N	<u>.</u>											
45	(vi)	ORIGINAL	. SOURCE												

	(A) ORCANIEM: Ecohorichia coli (B) STRAIN: K-12 RR1	
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:	
	CGCCGATTTA CCGCCTTGAG	20
10	(2) INFORMATION FOR SEQ ID NO:11:	
15	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOTHETICAL: NO	
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	
	CCGTCTGGAA CCACGCCCGA T	21
25	(2) INFORMATION FOR SEQ ID NO:12:	
30	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
35	(iii) HYPOTHETICAL: NO	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	
	ATCGGGCGTG ATTCCAGACG G	21
40		
	Claims	
45	 A microorganism transformed with a recombinant DNA sequence encoding an enzyme white responds to prephenate dehydratase in which one or two amino acid residue(s) are substitute other amino acid residue(s) or one or more amino acid residue(s) are deleted, and wherein fe 	ed wit

- corwith ack inhibition is released.
- 2. A microorganism as claimed in claim 1, wherein said microorganism is of the genus Escherichia, said prephenate dehydratase is chorismate mutase - prephenate dehydratase, derived from the genus 50 Escherichia and the serine residue 330 residues from the N-terminus is substituted with another amino acid residue.
- 3. A microorganism as claimed in claim 2, wherein said serine residue is substituted with a proline residue 55 or an aspartic acid residue.
 - 4. A microorganism as claimed in claim 1, wherein said microorganism is of the genus Escherichia, said prephenate dehydratase is chorismate mutase - prephenate dehydratase derived from the genus

Escherichia and the amino acid residues downstream from the 330 serine residue are deleted.

- A microorganism as claimed in claim 1, wherein said microorganism is a Coryneform bacterium, said prephenate dehydratase is derived from a Coryneform bacterium and the serine residue 235 residues from the N-terminus is substituted.
- 6. A microorganism belonging to the genus Escherichia transformed with a recombinant DNA sequence encoding an enzyme corresponding to 3-deoxy-D-arabinoheptulonic acid-7-phosphate synthase derived from the genus Escherichia, wherein one or two amino acid residue(s) are substituted and feedback inhibition is released.
- A microorganism as claimed in claim 6, further comprising a recombinant DNA sequence encoding one
 or more enzymes in the biosynthesis of an aromatic amino acid, wherein feedback inhibition of said
 enzyme is released.
- 8. A microorganism as claimed in claim 6 or 7, wherein said 3-deoxy-D-arabinoheptulonic acid-7-phosphate synthase is encoded by aroF.
- 9. A microorganism as claimed in claim 8, wherein the aspartic acid residue 147 residues from the N-terminus or the serine residue 181 residues from the N-terminus of said 3-deoxy-D-arabinoheptulonic acid-7-phosphate synthase is substituted with another amino acid residue.
 - 10. A microorganism as claimed in claim 9, wherein said aspartic acid residue is substituted with an asparagine residue.
 - A microorganism as claimed in claim 9, wherein said serine residue is substituted with a phenylalanine residue.
- **12.** A microorganism as claimed in claim 6 or 7, wherein said 3-deoxy-D-arabinoheptulonic acid-7-phosphate synthase is encoded by aroG.
 - 13. A microorganism as claimed in claim 12, wherein said one or two amino acid residue(s) of said 3-deoxy-D-arabinoheptulonic acid-7-phosphate synthase being substituted are selected from the group consisting of the aspartic acid residue 146 amino acid residues from the N-terminus, the methionine residue 147 amino acid residues from the N-terminus, the proline residue 150 amino acid residues from the N-terminus, and both the methionine residue 157 amino acid residues from the N-terminus and the alanine residue 219 amino acid residues from the N-terminus.
- 40 14. A microorganism as claimed in claim 13, wherein said aspartic acid residue 146 amino acid residues from the N-terminus is substituted with an asparagine residue, said methionine residue 147 amino acid residues from the N-terminus is substituted with an isoleucine residue, said proline residue 150 amino acid residues from the N-terminus is substituted with a leucine residue, said alanine residue 202 amino acid residues from the N-terminus is substituted with an isoleucine residue and said alanine residue 219 amino acid residues from the N-terminus is substituted with an threonine residue.
 - 15. A microorganism as claimed in claim 7, wherein said enzyme is chorismate mutase-prephenate dehydratase.
 - 16. A microorganism as claimed in claim 7, wherein said enzyme in the biosynthesis of an aromatic amino acid is a tryptophan operon.
- 17. A process for preparing an aromatic amino acid which comprises culturing a microorganism claimed in claims 1, 2, 4, 5, 6 or 7 in a medium, and isolating the aromatic amino acid produced.
 - 18. A process for preparing an aromatic amino acid as claimed in claim 17, wherein said aromatic amino acid is L-phenylalanine or L-tryptophan.

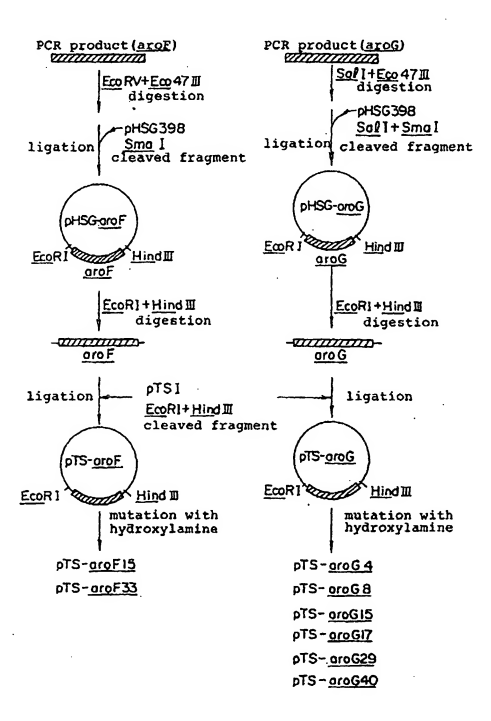
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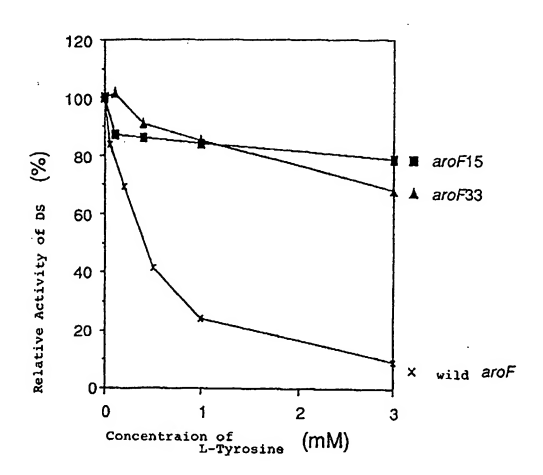
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35

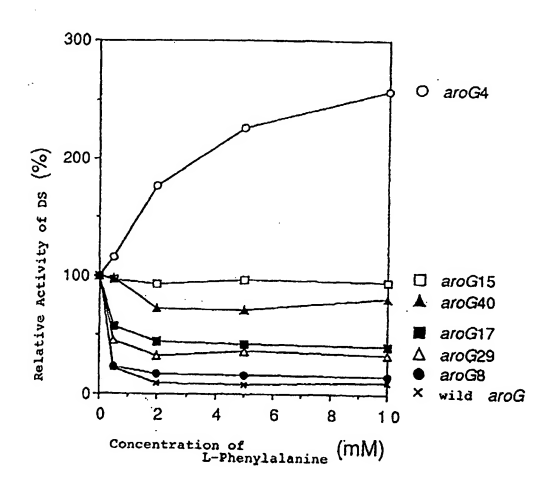
- 19. A recombinant DNA sequence encoding an enzyme which corresponds to prephenate dehydratase derived from a Coryneform bacterium, 3-deoxy-D-arabinoheptulonic acid-7-phosphate synthase, or chorismate mutase-prephenate dehydratase derived from the genus Escherichia, in which one or two amino acid residue(s) are substituted with other amino acid residue(s) or one or more amino acid residue(s) are deleted, and wherein feedback inhibition is released.
- 20. An enzyme which corresponds to prephenate dehydratase derived from a Coryneform bacterium, 3-deoxy-D-arabinoheptulonic acid-7-phosphate synthase, or chorismate mutase-prephenate dehydratase derived from the genus Escherichia, wherein one or two amino acid residue(s) are substituted or one or more amino acid residue(s) are deleted, and feedback inhibition is released.



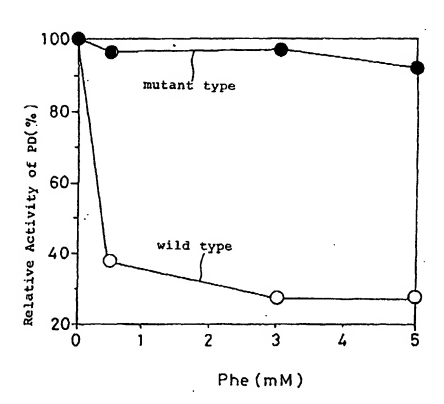




(FIG. 3)







(FIG. 5)

